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(54) Title: HISTIDINE-TAGGED SHIGA TOXINS, TOXOIDS, AND PROTEIN FUSIONS WITH SUCH TOXINS AND TOXOIDS, METHODS FOR THE PURIFICATION AND PREPARATION THEREOF

(57) Abstract

The present invention describes the isolation and purification of biologically and immunologically active histidine-tagged Shiga toxins (His-tagged), a toxin associated with HC and the potentially life-threatening sequela HUS transmitted by strains of pathogenic bacteria. The present invention describes how his-tagging greatly simplifies and expedites purifying Shiga toxins, and describes an improved method for such purification. One aspect of the invention is obtaining and using Shiga toxins or toxoids that are immunoreactive but not toxic. Another aspect of the invention is obtaining and using fusion proteins of His-tagged Shiga toxins or toxoids. Yet another aspect of the invention is obtaining and using antibodies to His-gagged Shiga toxins, toxoids, or Shiga toxin/toxoid fusion proteins.

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-1-

Description

HISTIDINE-TAGGED SHIGA TOXINS, TOXOIDS, AND PROTEIN FUSIONS WITH SUCH TOXINS AND TOXOIDS, METHODS FOR THE PURIFICATION AND PREPARATION THEREOF

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GOVERNMENT INTEREST

The invention described herein may be manufactured, licensed, and used for governmental purposes without payment of royalties to us thereon.

FIELD OF THE INVENTION

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The invention relates to a family of multi-unit bacterial proteins that are associated with hemorrhagic colitis and the life-threatening sequela, hemolytic uremic syndrome. These proteins, defined as members of the "Shiga toxin family," have been tagged with histidine residues. The invention further relates to a non-toxinogenic but immunoreactive form of histidine-tagged Shiga toxins, or toxoids. Moreover, the invention relates to fusion proteins obtained by combining histidine-tagged Shiga toxins or toxoids with other proteins. Histidine tagging greatly facilitates purification of Shiga toxins, and the invention also relates to methods for purifying such toxins. The invention further relates to using the histidine-tagged Shiga toxoids or fusion proteins of Shiga toxoids as antigens for generating an immune response against infection or transmission by bacteria expressing Shiga toxin. It also relates to antibodies to Shiga toxins, toxoids, or Shiga toxin/toxoid fusion proteins, both monoclonal and polyclonal, and their use in treating, diagnosing, and preventing of disease and infections by pathogenic *E. coli*. Finally, the invention relates to preparing the Shiga toxins, toxoids, and fusion proteins.

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BACKGROUND OF THE INVENTION

Enterohemorrhagic Escherichia coli (EHEC) are associated with food-borne outbreaks of bloody diarrhea or "hemorrhagic colitis" (HC) and the hemolytic uremic syndrome (HUS). (Spika, J. et al., "Hemolytic Uremic Syndrome and Diarrhea Associated with Escheria coli: 0157:H7 in a Day Care Center," J. Pediatr., 109: 287-291(1986); Remis, R., "Sporadic case of hemorrhagic colitis associated with Escheria coli 0157:H7," Ann. Intern. Med., 101:624-626 (1984); "Riley, L. et al., "Hemorrhagic

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colitis associated with a rare *Escheria coli* serotype," *N. Engl. J. Med.*, 308:681-685 (1983)). EHEC infection can be deadly and poses a significant threat to the young and the elderly, who are the most likely to develop serious complications from EHEC infections. Several outbreaks and sporadic cases of HC and HUS have occurred over the past few years, with the largest outbreak in United States in 1993. In that outbreak, over 500 cases of HC and HUS were traced to contaminated hamburgers from a Jack-in-the Box fast food restaurant. (Centers for Disease Control and Prevention, *Morbid. Mortal. Weekly Rep.*, 42:258(1993)). In July 1996, a large outbreak of EHEC in Japan resulted in over 10,000 infected individuals and 8 deaths. Many Japanese children required hospitalization. Unfortunately, no cure or vaccine for HC and HUS is currently available.

Primarily, HC and HUS are transmitted by the ingestion of contaminated food, particularly undercooked beef products, such as hamburger. (Doyle et al., *J. Appl. Environ. Microbiol.* 53:2394 (1987); Samadpour et al., *J. Appl. Environ. Microbiol.* 60:1038 (1994)). With the prevalence of EHEC in cattle and the subjective nature of differentiating between cooked and undercooked hamburgers, a stop at a fast food restaurant or a family barbecue can result in tragedy. HC and HUS appear to be mediated by the toxin produced by EHEC and *Shigella dysenteriae* (for review see O'Brien and Holmes, *Microbiol.* Rev., 51: 206-220 (1987)). These bacteria produce a family of closely related cytotoxins that collectively will be called "Shiga toxins" for the purpose of this application. Shiga toxins (alternatively, "verotoxins") have cytotoxic, neurotoxic, and enterotoxic activity (Strockbine, N. et al., "Two toxinconverting phages from *Escheria coli* 0157:H7 strain 933 encode antigenically distinct toxins with similar biological activities," *Infect. Immun.*, 53:135-140 (1986)).

Based on their immunological cross-reactivity, the Shiga toxins have been divided into two groups. (Strockbine et al., *supra*). These groups have been designated Shiga toxin type 1 (Stx1) and Shiga toxin type 2 (Stx2). (Strockbine et al., *supra*; Calderwood et al., "Proposed New Nomenclature for SLT (VT) Family," ASM News, 62:118-119 (1996)). The Stx1 group contains the prototype Stx1 toxin from

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EHEC as well as the Shiga toxin from *Shigella dysenteriae* type 1. In recent years, other types of toxins have been discovered and considered members of the Stx2 group. These are Stx2e, Stx2c, Stx2vha, and Stx2vhb. (Lindgren et al., *Infection and Immunology*, 61:3832 (1993); Schmitt, C. et al., "Two copies of Shiga-like toxin II-related genes common in enterohemorrhagic *Escheria coli* strains are responsible for the antigenic heterogeneity of the 0157:H strain E325II," *Infect. Immun.*, 59:1065-1073 (1991); Marques, L. et al., "*Escheria coli* strains isolated from pigs with edema disease produce a variant of Shiga-like toxin II," *FEMS Lett.*, 44:33-38 (1987)).

For the purposes of this application the term "Shiga toxin" encompasses Shiga toxin and any other toxins in the Stx1 or Stx2 group. The abbreviation "Stx" will refer to the protein designation, and the abbreviation "stx" to the gene designation.

These Shiga toxins do share similar genetic and protein organization, as set forth in Figure 1. The A subunit gene encodes the enzymatically active subunit. The A subunit polypeptide has two functional domains, A1 and A2, which are linked by a disulfide bond. The Al portion is an N-glycosidase that acts on the 28S rRNA subunit of eukaryotic ribosomes to inhibit protein synthesis. (Saxena, S. et al., "Shiga toxin, Shiga-like toxin II variant, and ricin are all single-site RNA N-glycosidases of 28S RNA when microinjected into *Xenopus* oocytes," *J. Biol. Chem.*, 264:596-601 (1989)). The A2 fragment is required for the binding of 5 B subunit polypeptides. The pentamer of B subunits is responsible for binding to a receptor on eukaryotic cells. A polypeptide containing the entire A subunit and B subunit pentameter is referred to as a Shiga holotoxin. Despite this knowledge about the toxin components, there is no known cure or vaccine for HC or HUS.

The need exists for therapeutic agents for the treatment and prevention of HC and HUS. However, progress in the search for such agents has been hampered by the lack of a fast and simple method for purifying Shiga toxins. Therefore, the need exists for such a fast and simple method. Moreover, the need exists for such a method that further allows for large-scale production of Shiga toxins while retaining their biological and immunological activity. The need also exists for such a method that

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-4-

allows for large-scale production of Shiga toxoids and fusion proteins of Shiga toxins and toxoids. Such a method should simplify obtaining antibodies against Shiga toxin and vaccines against HC and HUS using Shiga toxoids and fusions of Shiga toxoids.

SUMMARY OF THE INVENTION

The present invention describes the isolation and purification of biologically and immunologically active histidine-tagged Shiga toxins (His-tagged), a toxin associated with HC and the potentially life-threatening disease HUS transmitted by strains of pathogenic bacteria. The present invention describes how his-tagging greatly simplifies and expedites purifying Shiga toxins, and describes an improved method for such purification.

One aspect of the invention is obtaining and using Shiga toxoids that are immunoreactive but not toxic. For example, the invention describes using such obtained Shiga toxoids in vaccines against HC and HUS.

Another aspect of the invention is obtaining and using fusion proteins of Histagged Shiga toxins or toxoids. These fusion proteins have the advantage of combining beneficial properties of each protein, resulting, for example, in improved protein stability or targeted delivery of a his-tagged Shiga therapeutic agent.

Yet another aspect of the invention is obtaining and using antibodies to Histagged Shiga toxins, toxoids, or Shiga toxin/toxoid fusion proteins. These antibodies can be either monoclonal or polyclonal and have potential uses in treating, diagnosing, or preventing HC and HUS caused by EHEC or *Shigella dysenteriae* type 1 infections.

Other aspects of the present invention will become apparent from the more detailed description provided below, to be read in conjunction with the accompanying drawings.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 depicts the protein structure of Shiga toxin genes.

Figure 2 depicts the predicted amino acid sequence for the mature A subunit and the unprocessed B subunit of Stxl. (Calderwood et al., *Proc. Natl. Acad. Sci. USA*, 84: 4364-4368 (1987); DeGrandis et al., *J. Bacteriol.*, 169:4313-4319(1987)).

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Figure 3 depicts the predicted amino acid sequence for the mature A subunit and the unprocessed B subunit of Stx2. (Jackson et al., *FEMS Lett.*, 44:109-114 (1987)).

Figure 4 depicts the predicted DNA sequence for stx1 and DNA upstream of that sequence. (Calderwood et al., Proc. Natl. Acad. Sci. USA, 84: 4364-4368 (1987); DeGrandis et al., J. Bacteriol., 169: 4313-4319 (1987)).

Figure 5 depicts the predicted DNA sequence for stx2 and DNA upstream of that sequence. (Jackson et al., FEMS Lett., 44: 109-114 (1987)).

Figure 6 depicts the approximately 1200 base pair fragments of stx1 produced by PCR amplification. Figures 6a-c depict the fragments used to make plasmids pQHI, pQHEI, and p7HI, respectively. Nucleotides in lower case represent non-toxin sequences in the primers and/or base changes.

Figure 7 depicts the approximately 1200 base pair fragments of stx2 produced by PCR amplification. Figures 7a and 7b depict the fragments used to make plasmids pQHI and pQHEII, respectively. Nucleotides in lower case represent non-toxin sequences in the primers and/or base changes.

Figure 8 depicts the plasmid pQHI, encoding the His-Stx I fusion and driven by the T5 promoter.

Figure 9 depicts the plasmid pQHII, encoding the His-Stx 2 fusion and driven by the T5 promoter.

Figure 10 depicts the plasmid pQHEI, encoding the His-Enterokinase site-Stx1 fusion and driven by the T5 promoter.

Figure 11 depicts the plasmid pQHEII, encoding the His-Enterokinase site-Stx 2 fusion and driven by the T5 promoter.

Figure 12 depicts the plasmid pQHIIvhb, encoding the His-Stx 2 fusion and driven by the T5 promoter.

Figure 13 depicts the plasmid pQHEIIvhb, encoding the His-Enterokinase site-Stx 2 fusion and driven by the T5 promoter.

-6-

Figure 14 depicts the plasmid p7HI, encoding the His-Stx I fusion and driven by the PT7 promoter.

Figure 15 depicts the plasmid p7HII, encoding the His-Stx 2 fusion and driven by the PT7 promoter.

Figure 16 depicts the expression of His-Stx fusion proteins according to the invention.

DETAILED DESCRIPTION OF THE INVENTION

An object of the invention is to purify large quantities of Shiga toxins that retain their biological and immunological properties. To achieve this object, a Shiga toxin gene was cloned into a Histidine-tag expression vector, expressed, and purified. An additional object of the invention is to obtain antigens specific to Shiga toxoids, a toxin that is non-toxinogenic but immunoreactive, for generating an immune response against Shiga toxins. Another object of the invention is the creation of antibodies against Shiga toxins or toxoids for treating, diagnosing, or preventing disease and infections by pathogenic bacteria. The his-tagged Shiga toxins or toxoids described above can be used for these purposes.

Those skilled in the art will also recognize that the size of the his-tagged Shiga toxin to be used may be varied according to the specific purpose for the Shiga toxin. For example, if the purpose is fusing the his-tagged Shiga toxin or toxoid with one or more proteins, a smaller fragment might be selected to enhance stability of the combined fusion product, although using a larger fragment is by no means precluded. The desired size of the His-Shiga toxin may also vary with the convenience of the available restriction sites, in light of the materials and methods known to those skilled in the art. Consequently, the terms "His-Shiga toxin" or "His-tagged Shiga Toxin" refers to the fragment of about 372-377 amino acids comprising the A and B subunits of any of the Shiga toxin family members fused with a histidine tag. Smaller fragments that retain biological and/or immunological function are also included. Biological function is measured by, for example, cytotoxicity to Vero cells, as described in Example III.A. Immunological function may also be tested by, for

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-7-

example, neutralization by specific antisera, as described in Example III.B. A preferred embodiment of the invention is a His-Tagged Shiga holotoxin, containing 1 A subunit and 5 B subunits. In another preferred embodiment, the tag consists of six histidine residues. The most preferred embodiment is a His₆-Tagged Shiga holotoxin.

One of the objects of the present invention is to administer His-Shiga toxoids to protect against illness or disease caused by EHEC or *Shigella dysenteria* type I, such as HC and HUS. The object is achieved through the stimulation of immune response directed against Shiga toxins.

Consequently, the term "immunizing" or "immunization" is used in the application. The degree of protection achieved by such immunization will vary with the degree of homology between Shiga toxins and the His-Shiga toxoids, as well as other factors, such as unique attributes of the patient or the species treated. Moreover, immunization is not limited to avoiding infection altogether; it also includes decreasing the severity of the infection, as measured by the following indicators: reduced incidence of death, HUS, or permanent kidney damage; decreased levels of toxin; reduced fluid loss; or other indicators of illness regularly used by those skilled in the relevant art.

Unless specified otherwise, the uses and methods set forth herein are generally applicable to humans and animals. The term "patient" is used herein to mean both humans and animals, and "animals" is not limited to domesticated animals but also may include wildlife and laboratory animals.

Moreover, because His-Tagged Shiga toxins according to the invention have the biological and immunological properties of Shiga toxins, they may be used for any application appropriate for Shiga toxins. For example, it has been recently demonstrated that Stxl can be used to treat bone marrow cells from mice with human B-cell lymphomas. The Shiga toxin bound to the receptor on the lymphoma cell and the toxin killed the cancer cell. (LaCasse et al., *Blood* 88:1551(1996)). Thus, the skilled artisan would expect that His-Shiga toxins or fusions could be used for the same purpose and in the same manner.

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-8-

Isolating and Purifying His-Tagged Shiga Toxin

The standard protocol for purification of Shiga toxin comprising A and B subunits uses biochemical techniques. The standard protocol was developed by O'Brien et al. (O'Brien et al., *Infect. Immun.* 40:675 (1983); O'Brien et al., *Infect. Immun.*, 30:170(1980)). The method employs four purification steps: 1) ammonium sulfate precipitation; 2) DEAE Sepharose column chromotography; 3) chromatofocusing; and 4) antibody affinity chromotography. This method has the advantages of employing publicly available materials, being capable of purifying all Shiga toxins, and being capable of purifying Shiga toxins for human use. Its disadvantage is that the minimum time required for this test is three weeks.

Another well-known method for purifying Shiga toxin from bacteria was developed by Keusch et al. (Donohue-Rolfe et al., *Infect. Immun.* 57:3888 (1989); Acheson et al., *Microb. Pathog.* 14:57 (1993)). This method employs a Shiga toxin receptor analog. The receptor analog is the P1 glycoprotein (P1gp) from tapeworm hydatid cysts material (HCM) in sheep gut. This method contains three purification steps: 1) ammonium sulfate precipitation; 2) Blue scpharose chromotography; and 3) Plgp column chromatography. The Plgp must be prepared from the HCM. Though faster than the standard method, this method still requires a minimum of two or more weeks. The hydatid cyst material must be obtained from infected sheep and is not publicly available. The method has the additional disadvantages of being capable of use with only those Shiga toxins that bind P1gp and, because of possible contamination, it is not appropriate for obtaining Shiga toxoids for use in humans.

Using recombinant methods, the Shiga toxin gene or portions of the Shiga toxin gene have been cloned and expressed in bacteria and purified. Zollman et al., *Prot. Expression Pur.* 5:291 (1994), purified a recombinant Stx1 A1 fragment. Acheson et al., *Infect. Immunol.* 63:301(1995), expressed and purified the Stx2 B subunit. Downes et al., *Infect. Immun.* 56:1929 (1988), expressed the *stx2* gene in bacteria and purified Stx2. However, the purification methods following expression

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-9-

were essentially those of the standard method or hydatid cyst method and, therefore, had the same disadvantages.

In the search for a method for purifying Shiga toxins, applicants have developed a purification method based on the creation of a histidine-tagged Shiga toxin. Methods for histidine tagging are known in the art. For example, Fryxell et al., Biochem. Biophys. Res. Comm., 210:253-259 (1995), added a kemptide and histidine tag to the A chain of the eukaryotic toxin ricin, which was later associated with the B subunit. The ricin toxin differs from Shiga toxins in origin (prokaryotic v. eukaryotic) and structure of the B subunit (the ricin B subunit is a single polypeptide, not a pentamer). Moreover, Fryxell et al. only expressed the A subunit with a His-Tag. In addition, Strauss et al., FEMS Microbiol. Lett., 127:249-254 (1995), have his-tagged the C-terminus of the cholera toxin B subunit, and expressed a his-tagged B subunit-IgA protease fusion protein. However, this did not involve expressing the entire toxin with a His-Tag, and the expressed fusion protein did not undergo multimerization. Finally, Terbush & Novick, J. Cell. Biol., 130:299-312 (1995), tagged the C-terminus of a multiunit yeast protein. This involves a eukaryotic rather than a prokaryotic system. Moreover, expressing a functionally active Shiga toxin requires retaining its multimer conformation, as well as its receptor binding and enzymatic activity.

Although His-Tagging of proteins is known, it was not expected that His-Tagging of a Shiga toxin would be successful. The skilled artisan would have believed that a His-Shiga toxin fusion would have lost cytotoxicity, because the skilled artisan would have expected that the attachment of a His-Tag to the amino acid terminus of a toxin would destroy its activity. Moreover, the multi-unit toxin would have been expected to be more susceptible to losing toxicity upon fusion with additional amino acids, since it is known that the toxin must retain its conformation for enzymatic activity and for binding of the B subunits to cell receptors, and the addition of amino acids would have been expected to destroy proper conformation. The skilled artisan would be aware that conformation and the charge of the molecule is critical to Shiga toxins. For example, the skilled artisan would know that deleting a few N-terminal

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amino acids from Stx2A destroyed enzymatic activity, as reported in Perera et al., Infect. Immunol., 59:829-835 (1991)). Similarly, altering the C-terminus of the B subunit affected toxicity. (Perera et al., supra). Additionally, Perera et al. suggested that the charge of the molecule plays an important role. The importance of preserving Shiga toxin conformation is further underscored by findings that the highly homologous Stx1A and Stx2B subunits cannot be combined to form an active toxin. (Weinstein et al., Infect. Immun., 57:3743-3750 (1989)). Based on this knowledge, the skilled artisan would have expected that tagging a Shiga holotoxin with histidine residues would have unfavorably affected conformation and charge of the toxin product.

Surprisingly, his-tagging of Shiga toxin comprising A and B subunits generated a functional Shiga toxin, which has similar specific activity to Shiga toxin purified by standard methods. Moreover, the His-Shiga toxins are neutralized by monoclonal antibodies specific for Shiga toxins. Example I describes how to create the His-Shiga toxin fusion protein.

The following examples are intended to illustrate the invention but not to limit it. The skilled artisan will understand from these examples that modifications can be made that are still within the scope of the invention. The scope of the invention is defined by the claims.

20 Example I

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A. Construction of Plasmid Encoding His-Tagged Shiga Toxin

The His-Stx fusion clones were generated by PCR amplification of stx operons, restriction enzyme digestion of the PCR products, and ligation of the fragments inframe into the appropriate vectors. The expression vectors and primers were used to place histidine residues at the amino acid terminus of the toxins and place the constructs under the control of either an IPTG-inducible promoter (pQE vectors) (Qiagen, Inc., 9600 DeSoto Avenue, Chatsworth, CA 91311, 1-800-362-7737) or a T7 promoter (pt7-7) (Tabor et al., *Proc. Natl. Acad. Sci.* 82:1074 (1985)). The methods for obtaining His-Stx fusion clones are described in more detail below.

1. Bacterial strains and plasmids. The bacterial strains and plasmids used in this study are shown in Table 1.

Table 1. Bacterial strains and plasmids used in this study.

5	Strain or plasmid	Characteristic(s)	Source or reference
	E. coli strains		
	DH5α	Host strain for cloning	BRL
	XL1-Blue	Host strain for cloning; lacl; Tc1	Stratagene
	M15	Host strain for protein purification	Qiagen
	Plasmids		
	pJES120	Encodes stx II toxin operon	a
	рЈN25	Encodes stx 1 toxin operon	b
	pSQ543	Encodes stx IIvhb operon	c
	pQE30	Histidine fusion vector	Qiagen
	pQE32	Histidine fusion vector	Qiagen
	pREP4	lacl; Kn'	Qiagen
	pT7-7	T7 expression vector	d
	pGP1-2	Encodes T7 RNA polymerase; KN1	d

²⁰ a = Lindgren et al., Infect. Immunol. 61:3832 (1993).

2. Media and enzymes. Bacterial strains were grown in L broth (per liter: 10 g tryptone, 5 g yeast extract, 5 g NaCl). Kanamycin, tetracycline, and ampicillin (Sigma Chemical Co., St. Louis, MO.) were added to the medium at final concentrations of 25, 10, and 100 μg/ml (respectively) as needed. Restriction

b= Newland et al., "Cloning of shiga-like toxin structural genes from a phage of Escheria coli strain

^{933,} in Advances in Research on Cholera and Diarheas (S. Kuwahara & N.F. Pierce eds. 1994).

²⁵ c= Lindgren et al., Infect. Immunol. 62:623 (1994).

d= Tabor et al., Proc. Natl. Acad. Sci. USA 82:1074 (1985).

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endonucleases, calf intestinal phosphatase, and ligase were from Boehringer Mannheim, Indianapolis, Ind., or U.S. Biochemicals Corporation, Cleveland, Ohio. Enzymes were used according to manufacturer's instructions.

3. Primers and PCR. The Shiga toxin genes are cloned using polymerase chain reaction (PCR), a standard technique in the art. Primers were designed to amplify the stx toxin operons beginning at the first codon of the mature A subunit gene and ending downstream of the termination codon of the B subunit gene and created using standard techniques. The primers contained recognition sequences to generate unique restriction sites at the ends of the toxin operon. In a preferred embodiment, the 5' primers also contained sequences to encode the recognition sequence of the protease enterokinase to allow for removal of the histidine residues. The primers used are shown in Table 2.

TABLE 2. Primers used.

15	Primer	Primer Sequent (5'-3')	Restriction site
	HEC	GCGGATCCGATGACGATGACAAACGGGAGTTTACGATAGACTT	BamIII
	ПВАМ	GCGGATCCGGGAGTTTACGATAGACTT	BamHl
	11113	CCACGAATAAGCTTATGCCTCA	HindIII
20	IBAM5	GCGGATCCAAGGAATTTACCTTAGACTTC	BamHi
	IEC5	GCGGATCCGATGACGATGACAAAAAGGAATTTACCTTAGACTTC	Bamill
	IPST3	ATTTTCACTGCAGCTATTCTG	Pstl
	SI.TIIH5	GCATATGCATCACCATCACCATCACCGGGAGTTTACGATAGAC	Ndel
	SLTIH5	GCATATGCATCACCATCACCATCACAAGGAATTTACCTTAGECTTC	Ndel
25	SLT1.IC3	TAACATTTATCGATATCTCCGCCTG	Clal

Sequences encoding stx toxins were amplified from toxin clones using a PCR kit (GeneAmp kit, Perkin-Elmer Cetus, Norwalk, CT), which was used according to the manufacturer's instructions. The resulting stx PCR products were approximately

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1200 bp are shown in Figures 6a, 6b, 6c, 7a, and 7b. The DNA products contain the coding sequences for the mature A subunit and the unprocessed B subunit.

Procedures for cloning are well known in the art and are described in Maniatis, Molecular Cloning: A Laboratory Manual (1982)).

4. **DNA manipulations.** Plasmid DNA was isolated by the method of Holmes and Quigley, *Anal. Biochem.*, 114:193-197 (1981). Alternatively, plasmid DNA was purified using Qiagen columns (Qiagen Inc., Chatsworth, CA). PCR products were digested with restriction endonucleases and ligated into the pQE30/32 vectors (Qiagen, Inc.), or into the vector pT7-7. PCR reactions and ligations are summarized in Table 3.

TABLE 3: PCR Reactions and ligations.

	Plasmid	Primer			Resulting clone
	template	pair	Vector	Cloning sites	
;	pJES120	IIEC + IIH3	pQE30	BamHI/HindIII	pQHEII .
	pJES120	IIBAM + IIH3	pQE32	BamHI/HindIII	pQHII
	рЛN25	IEC5 + IPST3	pQE30	BamHI/PstI	pQHEI -
	pJN25	IBAM + IPST3	pQE30	BamHI/PstI	pQHI
	pJES120	SLTIIH5 + IIH3	pT7-7	Ndel/Hindlll	p7HII*
)	рЈN25	SLTIH5 + SLTIC3	pT 7- 7	Ndel/Clal	р7НІ
	pSQ543	IIEC + IIH3	pQE30	BamHI/HindIII	pQHEIIvhb ^a
	pSQ543	IIBAM + IIH3	pQE32	BamHI/HindIII	pQHIIvhb ^a

^a The construction of these plasmids is in progress.

To illustrate, the clone pQHEII was constructed as follows:

Plasmid pJES120 was the template with primers IIEC and IIH3 in a PCR reaction for amplification of the stx2 operon. The resulting PCR product started with the first codon of the mature A subunit gene, extended through the A subunit gene. the complete B subunit gene, and ended just downstream of the terminate codon of the B subunit gene (Figure 6b). The PCR product was digested with the restriction endonucleases Bam HI and Hind III, as was the vector plasmid pQE30. The vector

pQE30 was chosen because ligation of the PCR product with pQE30 at the BamHI sites would result in an in-frame protein fusion of the 6 histidine residues, the enterokinase cleavage site, additional amino acids, and the +1 residue of the mature A subunit. The digested PCR product was ligated into the digested vector pQE30. The ligation reaction was transformed into strain XL1-Blue and plated on agar that contained ampicillin. Colonies were screened for the presence of a plasmid that contained an approximately 1200 bp BamHI/HindIII DNA insert. Clones were confirmed by IPTG induction of toxin expression (Example II) with a subsequent test for cytoxicity on vero cells (Example III). Positive clones were then transformed into M15(pREP4) for large scale production of toxin.

Example II

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Large scale purification of His-Tagged Shiga Toxins

His-Shiga toxin was purified under nondenaturing conditions because of the multi-subunit nature of the Shiga toxins. The strain was streaked onto a selective agar plate and incubated at 37°C for 18-24 hrs. A 20 ml overnight culture was then prepared from a colony. The saturated culture was then diluted 1/50 into one liter of L broth with antibiotics and the culture was grown at 37°C until it reached an O.D.600 of 0.7-0.9. IPTG (2mM final concentration) was then added to induce expression of the His6 tagged toxin and the culture was grown for an additional 5 hrs. Cells were pelleted and the pellet was kept at -70°C overnight. The pellet was resuspended in sonication buffer (50 mM sodium phosphate (pH 8.0), 300 mM sodium chloride, 20 mM imidazole, 30 μg/ml PMSF), and the cells were sonicated to release toxin. Alternatively, the cells were treated with polymixin-B (2 mg/ml final concentration) for 3 hrs at 4°C. The extracts were clarified by centrifugation and filtered through a milipore 0.45 μm filter.

The nickel-nitrilotriacetic acid ligand (Ni-NTA) gel was equilibrated with sonication buffer and the cell extract was added to the gel. Protein was allowed to bind for 1 hr at room temperature or at 4°C. The gel was washed with sonication buffer followed by wash buffer (50 mM sodium phosphate (pH 8.0), 300 mM sodium

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chloride, 20 mM imidazole, 30 µg/ml PMSF, 10% glycerol, 1% tween-20). Protein was eluted from the gel with a gradient of imidazole (0-500 mM in wash buffer without tween-20), and 1 ml fractions were collected.

Fractions were tested for cytotoxicity on Vero cells (as explained in Example III) and were subjected to SDS-PAGE and silver stain. Fractions that were highly cytotoxic and relatively clean were pooled and dialyzed against sonication buffer. This pool was then placed onto a Ni-NTA spin column (Qiagen) to further purify the His6toxin and the resulting two fractions were dialyzed against PBS. A final cytotoxicity assay and BCA protein assay were performed for the determination of the specific activity of the purified toxin.

The protocol described above is modification of the non-denaturing protocol described by Qiagen to purify His-tagged proteins. However, the toxin that eluted contained many contaminants. To achieve purer His-Shiga toxin, modifications were made. Specifically, Tween-20 was added to the wash buffer, and the pH of the wash buffer was adjusted to 8. Also, a final Ni-NTA spin column was added.

This one-step His-affinity method for purifying His-Shiga toxin by an Ni-NTA column has several advantages over existing methods, as summarized in Table 4.

TABLE 4. Comparison of Toxin Purification Techniques

20	Purification Method	Minimum time required ^a	Stepsb	Materials available	Use for all Shiga toxins
	Standard	3 weeks	4	yes	yes
	Hydatid Cyst	2 weeks +	3	no	no
	His ₆ affinity	1 week	2	yes	yes

^a Time from streaking the strain onto an agar plate. This does not include the preparation of Plgp from hydatid cyst material which takes a minimum of 1.5 weeks. ^b This does not include the multiple steps involved in the purification of Plgp from hydatid cyst material and preparation of the column.

The Ni-NTA one-step method is superior because of its relative speed and simplicity. It requires a minimum of one week as opposed to a minimum of two or

-16-

more weeks. Moreover, all of the materials are readily available, the method is not limited to Shiga toxins that bind P1gp, and the products are suitable for use in humans.

The Shiga toxin obtained by the method has many uses. For example, the His-Shiga toxin may be used as a positive control antigen in a Shiga toxin detection kit. Such kits will use a purified His-Shiga toxin as positive indicator for the toxin in a sample. Other uses are detailed in the Examples below.

Example III

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Verifying Biological and Immunological Activity of His-Shiga Toxins

A. Vero Cell Cytotoxicity Assay

The cytotoxicity of His-Shiga toxins obtained according to the methods described in Examples I and II was verified by determining their cytotoxicity for Vero cells. Cytotoxicity assays on strains that expressed His-Shiga toxins were done essentially as described by Gentry and Dalrymple, *J. Clin. Microbiol*, 12: 361-366 (1980). Briefly, cultures induced for the expression of His-Shiga toxins were disrupted by sonic lysis and clarified by centrifugation. The extracts were serially diluted in tissue culture medium (Dulbecco modified Eagle medium containing 10% fetal calf serum, 0.8 mM glutamine, 500 U of penicillin G per ml, and 500 mg of streptomycin per ml). One hundred microliters of 10-fold dilutions of the lysates were added to microtiter plate wells containing about 10⁴ Vero cells in 100 μl of medium. The tissue culture cells were incubated at 37°C in 5% CO₂ for 48 hours and then fixed and stained with crystal violet. The intensity of color of the fixed and stained cells was measured with a Titertek reader at 620 nm.

B. Antisera Neutralization Assay

His-Shiga toxins obtained according to the methods described in Examples I and II were tested for antisera neutralization. Neutralization of cytoxic activity was described in great detail in Schmitt et al., *Infect. and Immun.*, 59:1065-1073 (1991). Briefly, lysates were incubated with serial dilutions of monoclonal or polyclonal antisera specific for Stxl or Stx2 at 37°C for 2 hours. One hundred microliters of the

-17-

samples were then added to vero cells as described above. Percent neutralization was determined by the following formula:

 $\{[A_{620}(toxin + antibody) - A_{620}(toxin)]/A_{620}(untreated cells)\} \times 100.$

Example IV

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Constructing Fusions with His-Shiga Toxins and Other Proteins

Using methods well-known in the art, the His-Shiga toxin could be fused with another protein of interest. These methods include chemical and genetic methods, as in cloning and expressing a fusion protein, although one skilled other methods are readily apparent to one skilled in the art. (D.V. Goeddell, *Meth. Enzymol.* Vol. 185(1990); Itakura, *Science* 198:1056 (1977)). For example, if a combination vaccine for immunization against Shiga toxin and another toxin (protein X) is desired, then these two toxins can be fused into a single protein. This can be achieved by first cloning the codons for the histidine residues in frame to the coding region of protein X. The fragment containing His-Protein X is then subcloned in-frame of the Shiga toxin operon. In a preferred embodiment, the fragment is subcloned in-frame to the A2-B portions of the Shiga toxin operon. The resulting His-Protein X-A2-B5 fusion would ideally result in immunization against Shiga toxin and protein X.

One skilled in the art would recognize that various proteins from pathogens and haptens may be conjugated to a His-Shiga toxin. Haptens and antigens may derive from but are not limited to bacteria, rickettsiae, fungi, viruses, parasites, drugs, or chemicals. They may include, for example, small molecules such as peptides, oligosaccarides, and toxins. Certain antimicrobial drugs, chemotherapeutic drugs having the capacity of being absorbed into the intestine may also be coupled to Shiga toxin for targeted delivery, since the B subunit pentamer binds to receptors in the intestine. Conjugation methods are well known in the art. Exemplary methods are set forth in Goeddel, "Systems for Heterologous Gene Expression," *Meth. Ezymol.*, 185 (1990), Itakura, "Expression in *E. coli* of a chemically synthesized gene for the hormone somatostatin," *Science*, 198:1056-1063 (1977), and Goeddel et al.,

"Expression of chemically synthesized genes for human insulin," *Proc. Natl. Acad. Sci.* USA, 281: 544-548(1979).

Conjugation may be achieved by genetically fusing His-Shiga toxoids by standard molecular techniques or by conjugation to a polysaccharide. Methods of conjugation include those outlined in M. Brunswick et al., *J. Immunol.*, 140:3364 (1988) and Chemistry of Protein Conjugates and Crosslinking, *CRC Press*, Boston (1991). Coupling of Shiga toxids to other proteins or polysaccharides would prevent disease from additional pathogens.

Example V

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10 His-Shiga Toxoids

A form of Shiga toxin that is immunoreactive but not toxinogenic is needed for immunization in animals. Such a His-Shiga toxoid can be generated using chemical or genetic methods. The chemical method involves treating the His-Shiga toxin with either formaldehyde or glutaraldehyde, as described by Perera et al., *J Clin. Microbiol.* 26:2127 (1988)). Briefly, samples of toxin containing 100 μg of protein are treated for 3 days at 37°C with 0.1 M Na₂HPO₄ (pH 8.0) containing 1% formaldehyde, and the residual formaldehyde is removed by dialysis against phosphate-buffered saline (PBS). To prepare His-Shiga toxoid by treatment with glutaraldehyde, crude toxin samples containing 50 μg of protein are incubated at 37°C in 0.11% glutaraldehyde in 0.1 M Na₂HPO₄ (pH 8.0) for 30 min. The toxoid is then tested on Vero cells, as described in Example III, for loss of cytotoxicity.

Genetically, a toxoid may be produced by site-directed mutagenesis, as described in Gordon et al., *Infect. Immun.* 60:485 (1992); Hovde et al., *Proc. Natl. Acad. Sci.* 85:2568 (1988); Jackson et al., *J. Bacteriol.* 172: 3346-3350 (1990). Several methods and kits exist for site-directed mutagenesis of a gene. One method employs the Bio-Rad Muta-Gene *in vitro* mutagenesis kit. Oligonucleotides can be designed and synthesized which alter specific condons in the toxin genes. Uracilincorporated, single-stranded target plasmid DNA will be mutagenized according to

-19-

the directions supplied by the manufacturer of the mutagenesis kit. The nucleotide changes are then confirmed by DNA sequence analysis.

In the His-shiga toxin, two or more amino acids essential for enzymatic activity should be altered. For example, A subunit targets are the residues E167 and E170. The Shiga toxoid resulting from this mutation has been used for vaccinating pigs. (Gordon et al., *supra*).

Example VI

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Passive Immunity to Shiga Toxin Using His-Shiga Toxin and Toxoid

A. Antisera Specific for His-Shiga Toxins

Antisera specific for Shiga toxins are required to treat and prevent potentially-deadly infections by EHEC and *Shigella dysenteriae* type I. Specifically, once a child becomes infected, the child and his or her family members or children in his or her day care group can receive anti His-Shiga toxin sera to achieve a protective immune response. A protective immune response is one that elicits sufficient antibody to permit a patient to avoid infection, decrease the significance or severity of an infection, or decrease the ability of bacteria to colonize the gastrointestinal tract.

Animal studies have shown that administering anti-Shiga toxin sera to mice results in resistance to normally lethal infection of EHEC. (Lindgren et al., *Infect. Immun.* 62:623(1994); Wadolowski et al., *Infect. Immun.* 58:3959). Thus, applicants believe that administering anti-Shiga toxin sera to humans and other mammals would result in a protective immune response against Shiga toxin infections.

Methods are well-known in the art for producing antisera for passive immunization. For example, His-Shiga toxoid, obtained by the methods described in Example VI, can be administered to a mammal, such as a horse intraperitoneally. Currently, the horse is used to produce serum against botulism toxin for administration to humans, Hibbs et al., *Clin. Infect Dis.*, 23:337-40 (1996), and the horse would be a preferred method for producing shiga toxin antiserum. After several boosts with His-Shiga toxoid, the serum of the immunized horse (or other mammal) would be tested for neutralizing the cytotoxicity of Shiga toxins. Advantageously, a large amount of

serum can be quickly made using this method. However, patients must first be screened for an immune reaction to horse serum. For this purpose, a small amount of horse serum would be subcutaneously injected, and the patient would be monitored for a reaction. Such methods for administering horse antiserum against toxins to humans are well known to the skilled artisan. Hibbs et al., *supra*; Dehesa and Possani, *Toxicon*, 32: 1015-1018(1994); Gilan et al., *Toxicon*, 27:1105-1112 (1989).

More preferably, the His-Shiga toxoid can be administered to human volunteers, either intraperitoneally or orally. The plasma from these volunteers is then isolated, and the human anti-His-Shiga toxin serum can be administered to patients. No threat of serum sickness arises from this method. Human hyperimmune globulin to Hemophilus influenzae b, Streptococcus pneumoniae, and Neisseria meningitidis has previously been prepared by others (Siber et al., *Infect. and Immun.*, 45: 248-254 (1984)).

B. Vaccines Against Shiga Toxins

An embodiment of the invention is vaccines against Shiga toxin infection. For example, these vaccines can include antibodies directed against His-Shiga toxin, obtained further described in Example VII. Moreover, these vaccines can be combination vaccines that comprise His-Shiga toxoid fused or conjugated with another protein, hapten, or antigen, as described in Example IV. These vaccines can be administered intraperitoneally or injectably by methods well known in the art.

A preferred method of administering His-Shiga toxin or toxoid and fusions thereof is by further conjugation to Synsorb® (SynSorb Biotech, Inc., 1204 Kensington Road, N.W., Calgary, Alberta, Canada, T2N3P5.) Synosorb is a sand-like material to which Shiga toxin receptor (Gb3) is covalently bound (Armstrong et al., *J. Infect. Dis.*, 171:1042 (1995)). This compound has been shown to bind Shiga toxins and appears to be safe for human ingestion. (Armstrong et al., *supra*) The Synsorb is bound to the B subunit pentamer via the B subunit pentamer-receptor reaction. Conjugation with Synsorb adds further stability.

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Another embodiment of the invention involves the administration of nucleic acid vaccines. DNA encoding a His-Shiga toxoid is injected into a patient as naked DNA, or the DNA is delivered to the body by a carrier system such as retro viruses, adenoviruses, or other carriers known in the art. Following administration, the patient mounts an immune response against transiently expressed foreign antigens.

Currently nucleic acid vaccines, in general, are all nearing clinical trials. This approach to vaccines involves delivering the DNA encoding the desired antigen into the host by inserting the gene into a nonreplicating plasmid vector (Marwick, C. JAMA 273:1403(1995); reviewed in Vogel, F.R. and N. Sarver, *Clin. Microbiol.*, Rev. 8:406 (1995)).

The first published demonstration of the protective efficacy of such a vaccine has shown that intramuscular injection of plasmid DNA encoding influenza A virus (A/PR/8/34) nucleoprotein (NP) elicited protective immune responses in BALB/c mice against a heterologous strain of influenza virus (A/HK/68) (Ulmer, J.B. et al. *Science* 259:1745(1993)). Immunized animals had reduced virus titers in their lungs, decreased weight loss, and increased survival compared with challenged control mice. Both NP-specific cytotoxic T lymphocytes (CTL's) and NP antibodies were generated. The NP antibodies were ineffective at conferring protection, but the CTL's killed virus-infected cells and cells pulsed with the appropriate major histocompatibility complex class I-restricted peptide epitope.

Another study has shown that intramuscular injection of plasmid DNA encoding influenza virus A/PR/8/34 hemagglutinin resulted in the generation of neutralizing antibodies that protected mice against a heterologous lethal influenza virus challenge (Montgomery, D.L. et al. *DNA Cell Biol.*, 12:777 (1993)).

25 Example VII

WO 98/11229

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His-Shiga Antibodies

His-Shiga antibodies, polyclonal and monoclonal, can also be used in the treatment, diagnosis, and prevention of infections related to Shiga toxins. Because of their increased specificity, monoclonal antibodies are preferred. His-Shiga toxin

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antibodies can be administered to humans or other mammals to achieve a protective immune response, for treatment or prophylaxis. Antibodies, in a physiologically acceptable carrier, may be administered orally or intraperitoneally. For this purpose, monoclonal antibodies are preferred and humanized monoclonal antibodies are particularly preferred. Positive clinical responses in humans have been obtained with monoclonal antibodies, and one skilled in the art would know how to employ Shiga monoclonal antibodies in humans. See Fagerberg et al., "Tumor Regression in Monoclonal Antibody-treated Patients Correlates with the Presence of Anti-idiotype-reactive T Lymphocytes," Cancer Research, 55:1824-27 (1995); "A Phase I Study of Human/Mouse Chimeric Anti-ganglioside GD2 Antibody ch14.18 in Patients with Neuroblastoma," Eur. J. Cancer, 2:261-267 (1995)).

Another embodiment of the invention involves using antibodies to diagnose Shiga toxin infections. The antibody, using well-known methods of immunoassaying, is brought into contact with a sample from a patient, such as a fecal sample. In addition, the antibody may be used to detect Shiga toxins in sample taken from cow, such as cow feces. Moreover, meat may be tested using the anti His-Shiga toxin antibody for detection. A detection kit comprising the His-Shiga toxin antibody can be used for this purpose.

For example, a sandwich Elisa can be used. In this kit, rabbit anti-His-Shiga toxin antibody can be used to capture toxin from a sample to be tested. Goat anti-His-Shiga toxin antibody can then be added followed by a secondary antibody such as mouse α -goat antibody conjugated to horseradish peroxidase. The antibody can be detected by standard methods.

His-Shiga toxin polyclonal antibodies and monoclonal antibodies are described below.

A. Making Polyclonal Antibodies

The technique of Harlow, E. and D. Lane (eds.), *Antibodies- a Laboratory Manual*, Cold Spring Harbor, New York (1988), may be followed. The general procedure is outlined herein. Take pre-bleeds of each mouse to be immunized: Bleed

from the tail vein into an eppendorf tube. Incubate at 37°C for 30 min, stir gently with a sterile toothpick (to loosen the clot), store overnight at 4°C. In the morning, spin 10 min/10,000 rpm in the microfuge, and collect the serum (i.e., supernatant; red blood cells are the pellet). Store the serum at -20°C. The sera obtained will be used as a negative control after the mice are immunized.

Inject a BALB/c mouse intraperitoneally with 25 µg of His-Shiga toxoid (using Titremax adjuvant, according to the instructions of the manufacturer (CytRyx Corp., 154 Technology Pkwy., Norcross, GA. 30092, 800-345-2987)). Wait 2 weeks, boost with an identical shot, wait 7 days and bleed from the tail vein into an eppendorf tube. Incubate at 37°C for 30 min, stir gently with a sterile toothpick (to loosen the clot), store overnight at 4°C. In the morning, spin 10 min/10,000 rpm in the microfuge, and collect the serum. Store the sera at -20°C.

B. ELISA to test titer of Abs.

WO 98/11229

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The technique of Harlow, E. and D. Lane (eds.), *Antibodies: A Laboratory*15 *Manual*, Cold Spring Harbor, New York (1988), may be followed. The general procedure is outlined below:

- (1) bind His-Shiga toxoid to plastic microtiter plates at 50 ng/well in PBS. Incubate 2h/RT (room temp) or overnight at 4°C.
- (2) wash plate 2X with PBS.
- 20 (3) block wells with 100 μl blocking solution [3% bovine serum albumin (Sigma Chemical, St. Louis, MO.), 0.02% sodium azide (Sigma) in PBS store stock at 4°C] for l 2 h at RT.
 - (4) wash plate 2X with PBS.
- (5) primary Ab = 50 μl test sera diluted in blocking solution for example, start with
 1:50 and do eleven 1:2 dilutions, or start with 1:50 and do eleven 1:10 dilutions),
 incubate 2 h/RT.
 - (6) wash 4X with PBS.
 - (7) secondary Ab = goat horseradish-conjugated anti-mouse lg, affinity purified (Boehringer Mannheim Corp., 9115 Hague Rd., P.O. Box 50414, Indianapolis, IN.

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46250,800-262-1640). Add secondary Ab diluted 1:500 in blocking solution without azide. Incubate 1 h/RT.

- (8) wash 4X with PBS.
- (9) add 100 μl TMB Peroxidase substrate to each well (prepared according to the instructions of the manufacturer, BioRad Labs, 3300 Regatta Blvd., Richmond, CA. 94804). Allow blue color to develop (no more than 10 min). Stop the reaction with 100 μ1 H₂SO₄. Read the plate at 450 nm.

A titer is defined as an absorbance value ≥ 0.2 units above that obtained for mouse preimmune sera.

Anti-Shiga toxin Abs obtained from animals may be used clinically if one changes the specificity of the antibody to human. Such techniques are well known to those of ordinary skill in the art. G. Winter et al., "Man-made antibodies," *Nature*, 349: 293-299 (1991); P.T. Jones et al., "Replacing the complementarity-determining regions in a human antibody with those from a mouse," *Nature*, 321: 522-525 (1986); P. Carter et al., "Humanization of an anti-p185^{HER2} antibody for human cancer therapy," *Proc. Natl. Acad Sci. USA*, 89: 4285-4289 (1992). Such antibodies may be given to the sibling of an infected patient to reduce the risk of infection of the sibling.

C. Raising Monoclonal Antibodies to His-Shiga Toxin

Monoclonal antibodies directed against Shiga toxin are used to passively protect a patient against EHEC and Shigella dysenteriae type I infections. Monoclonal antibodies are generated from mouse cells, and the specificity of these antibodies are changed for use in humans. G. Winter et al., "Man-made antibodies," Nature, 349: 293-299 (1991); P.T. Jones et al., "Replacing the complementarity-determining regions in a human antibody with those from a mouse," Nature, 321:

522-525 (1986); P. Carter et al., "Humanization of an anti-pl85^{HER2} antibody for human cancer therapy," *Proc. Natl. Acad Sci. USA*, 89: 4285-4289 (1992). Monoclonal Abs represent a more "pure" antibody for administration to a patient.

The procedure outlined in Harlow, E. and D. Lane, Antibodies: A Laboratory Manual, Cold Spring Harbor, New York (1988) is followed: Five 4- to 5-week old

-25-

female BALB/cJ mice are prebled, and immunized intraperitoneally with 25 μg His-Shiga toxoid suspended in 100 μl of TiterMax. Mice are boosted twice in two week intervals, intraperitoneally with 25 μg His-Shiga toxoid suspended in 100 μl of TiterMax. Seven days after each boost, blood (~300 - 500 μl) is collected from the tail vein. Sera are assayed for the presence of anti-Shiga toxin antibody by ELISA (as described above).

Mice producing high titers of anti-His Shiga toxin antibodies are boosted both intravenously and intraperitoneally with 25 μg of His-Shiga toxoid in 100 μl of PBS, sacrificed three days later, and sera collected. Spleen cells are isolated and fused to Sp2/0-Ag mouse myeloma cells (ATCC #CRL1581) at a ratio of 10 spleen cells to 1 myeloma cell. Fused cells are distributed into microdilution plates, and culture supernatants are assayed by ELISA after 3-4 weeks of culture for anti-His-Shiga toxin antibodies. Cultures positive for production of anti-His Shiga toxin antibodies are expanded and cloned twice by limiting dilution.

The person skilled in the art would understand how to use and practice the invention based on the above disclosure. Other embodiments of the invention will be apparent to those skilled in the art from consideration of the specification and practice of the invention disclosed herein. It is intended that the specification and examples be considered as exemplary only, with the true scope and spirit of the invention being indicated by the following claims.

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We claim:

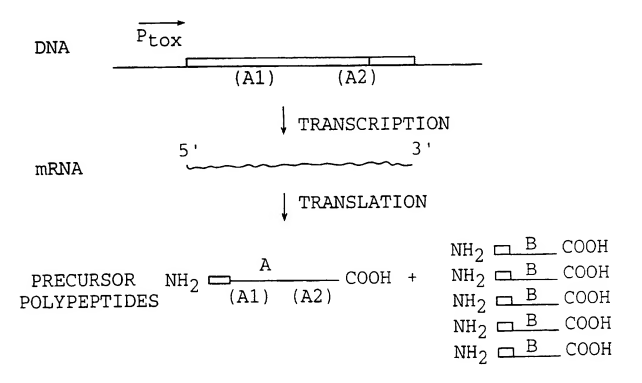
- 1. A polypeptide comprising a Shiga toxin having a histidine tag.
- 2. A polypeptide comprising an immunoreactive but non-toxinogenic form of the polypeptide of claim 1.
- 3. A fusion protein comprising the polypeptide of claim 1 or 2 fused to a second polypeptide or a portion thereof.
- 4. A method for large-scale isolation and purification of Shiga toxin comprising the steps of:
 - a) expressing Shiga toxin with a histidine tag in bacteria; and
 - b) eluting cell extract containing histidine-tagged Shiga toxin
- 5 over a nickel-

nitrilotriacetic acid ligand (Ni-NTA) gel.

- 5. A method of providing passive immune protection comprising the step of administering antisera directed against the polypeptide of claim 2 to patients in need thereof.
- 6. A method of treating infections mediated by toxins of the Shiga toxin family comprising the step of administering antibodies against the polypeptide of claim 2 to patients in need thereof.
- 7. A vaccine comprising an antibody directed against the polypeptide of claim 2.
- 8. A vaccine comprising a nucleotide encoding the polypeptide of claim 2.

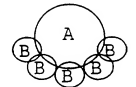
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1/19



PROCESSING, SECRETION, AND ASSEMBLY

MATURE TOXIN



P, PROMOTER. . , SIGNAL SEQUENCE.

FIG. 1

Stx 1A

1	KEFTLDFSTA	KTYVDSLNVI	RSAIGTPLQT	ISSGGTSLLM	IDSGSGDNLF
51	AVDVRGIDPE	EGRFNNLRLI	VERNNLYVTG	FVNRTNNVFY	RFADFSHVTF
101	PGTTAVTLSG	DSSYTTLQRV	AGISRTGMQI	NRHSLTTSYL	DLMSHSGTSL
151	TQSVARAMLR	FVTVTAEALR	FRQIQRGFRT	TLDDLSGRSY	VMTAEDVDLT
201	LNWGRLSSVL	PDYHGQDSVR	VGRISFGSIN	AILGSVALIL	NCHHHASRVA
251	RMASDEFPSM	CPADGRVRGI	THNKILWDSS	TLGAILMRRT	ISS*

Stx 1B

1	MKKTLLIAAS	LSFFSASALA	TPDCVTGKVE	YTKYNDDDTF	TVKVGDKELF
51	TNRWNLQSLL	LSAQITGMTV	TIKTNACHNG	GGFSEVIFR*	

FIG. 2

3/19

Stx 2A

1	REFTIDFSTQ	QSYVSSLNSI	RTEISTPLEH	ISQGTTSVSV	INHTPPGSYF
51	AVDIRGLDVY	QARFDHLRLI	IEQNNLYVAG	FVNTATNTFY	RFSDFTHISV
101	PGVTTVSMTT	DSSYTTLQRV	AALERSGMQI	SRHSLVSSYL	ALMEFSGNTM
151	TRDASRAVLR	FVTVTAEALR	FRQIQREFRQ	ALSETAPVYT	MTPGDVDLTL
201	NWGRISNVLP	EYRGEDGVRV	GRISFNNISA	ILGTVAVILN	CHHQGARSVR
251	AVNEESQPEC	QITGDRPVIK	INNTLWESNT	AAAFLNRKSQ	FLYTTG

Stx 2B

- 1 MKKMFMAVLF ALASVNAMAA DCAKGKIEFS KYNEDDTFTV KVDGKEYWTS
- 51 RWNLQPLLQS AQLTGMTVTI KSSTCESGSG FAEVQFNND*

FIG. 3

1	CTTGACCAGATATGTTAAGGTTGCAGCTCTCTTTGAATATGATTATCATTTTCATTACGT	bU
61	TATTGTTACGTTTATCCGGTGCGCCGTAAAACGCCGTCCTTCAGGGCGTGGAGGATGTCA	120
121	AGAATATAGTTATCGTATGGTGCTCAAGGAGTATTGTGTAATATGAAAATAATTATTTTT	180
181	AGAGTGCTAACTTTTTTCTTTGTTATCTTTTCAGTTAATGTGGTGGCGAAGGAATTTACC	240
241	TTAGACTTCTCGACTGCAAAGACGTATGTAGATTCGCTGAATGTCATTCGCTCTGCAATA	300
301	GGTACTCCATTACAGACTATTTCATCAGGAGGTACGTCTTTACTGATGATTGAT	360
361	TCAGGGGATAATTTGTTTGCAGTTGATGTCAGAGGGATAGATCCAGAGGAAGGGCGGTTT	420
421	AATAATCTACGGCTTATTGTTGAACGAAATAATTTATATGTGACAGGATTTGTTAACAGG	480
481	ACAAATAATGTTTTTTATCGCTTTGCTGATTTTTCACATGTTACCTTTCCAGGTACAACA	540
541	GCGGTTACATTGTCTGGTGACAGTAGCTATACCACGTTACAGCGTGTTGCAGGGATCAGT	600
601	CGTACGGGGATGCAGATAAATCGCCATTCGTTGACTACTTCTTATCTGGATTTAATGTCG	660
661	CATAGTGGAACCTCACTGACGCAGTCTGTGGCAAGAGCGATGTTACGGTTTGTTACTGTG	720
721	ACAGCTGAAGCTTTACGTTTTCGGCAAATACAGAGGGGATTTCGTACAACACTGGATGAT	780
781	$\tt CTCAGTGGGCGTTCTTATGTAATGACTGCTGAAGATGTTGATCTTACATTGAACTGGGGA$	840
841	${\tt AGGTTGAGTAGCGTCCTGACTATCATGGACAAGACTCTGTTCGTGTAGGAAGAATT}$	900
901	TCTTTTGGAAGCATTAATGCAATTCTGGGAAGCGTGGCATTAATACTGAATTGTCATCAT	960
961	CATGCATCGCGAGTTGCCAGAATGGCATCTGATGAGTTTCCTTCTATGTGTCCGGCAGAT	1020
1021	GGAAGAGTCCGTGGGATTACGCACAATAAAATATTGTGGGATTCATCCACTCTGGGGGCA	1080
1081	ATTCTGATGCGCAGAACTATTAGCAGTTGAGGGGGTAAAATGAAAAAAACATTATTAATA	1140
1141	${\tt GCTGCATCGCTTTCATTTTTTCAGCAAGTGCGCTGGCGACGCCTGATTGTGTAACTGGA}$	1200
1201	AAGGTGGAGTATACAAAATATAATGATGACGATACCTTTACAGTTAAAGTGGGTGATAAA	1260
1261	GAATTATTTACCAACAGATGGAATCTTCAGTCTCTTCTTCTCAGTGCGCAAATTACGGGG	1320
1321	${\tt ATGACTGTAACCATTAAAACTAATGCCTGTCATAATGGAGGGGGGATTCAGCGAAGTTATT}$	1380
1381	TTTCGTTGA 1389	

FIG. 4

1	ATCGCATAGCTCATCGGAACAAGCTCAAGCGGTCTCCGGTCGAGTCCTCATGCGTCCAT*!	60
61	ATCTGCATTATGCGTTGTTAGCTCAGCCGGACAGAGCAATTGCCTTCTGAGCAATCGGTC	120
121	ACTGGTTCGAATCCAGTACAACGCGCCATATTTATTTACCAGGCTCGCTTTTGCGGGCCT	180
181	$\verb"TTTTATATCTGCGCCGGGTCTGGTGCTGATTACTTCAGCCAAAAGGAACACCTGTATAT"$	240
241	GAAGTGTATATTTAAATGGGTACTGTGCCTGTTACTGGGTTTTTCTTCGGTATCCTA	300
301	TTCCCGGGAGTTTACGATAGACTTTTCGACCCAACAAGTTATGTCTCTTCGTTAAATAG	360
361	TATACGGACAGAGATATCGACCCCTCTTGAACATATATCTCAGGGGACCACATCGGTGTC	420
421	TGTTATTAACCACACCCCACCGGGCAGTTATTTTGCTGTGGATATACGAGGGCTTGATGT	480
481	$\tt CTATCAGGCGCGTTTTGACCATCTTCGTCTGATTATTGAGCAAAATAATTTATATGTGGC$	540
541	CGGGTTCGTTAATACGGCAACAAATACTTTCTACCGTTTTTCAGATTTTACACATATATC	600
601	AGTGCCCGGTGTGACAACGGTTTCCATGACAACGGACAGCAGTTATACCACTCTGCAACG	660
661	TGTCGCAGCGCTGGAACGTTCCGGAATGCAAATCAGTCGTCACTCAC	720
721	${\tt TCTGGCGTTAATGGAGTTCAGTGGTAATACAATGACCAGAGATGCATCCAGAGCAGTTCT}$	780
781	GCGTTTTGTCACTGTCACAGCAGAAGCCTTACGCTTCAGGCAGATACAGAGAGAATTTCG	840
841	TCAGGCACTGTCTGAAACTGCTCCTGTGTATACGATGACGCCGGGAGACGTGGACCTCAC	900
901	TCTGAACTGGGGGCGAATCAGCAATGTGCTTCCGGAGTATCGGGGAGAGGATGGTGTCAG	960
961	${\tt AGTGGGGAGAATATCCTTTAATAATATATCAGCGATACTGGGGACTGTGGCCGTTATACT}$	1020
1021	GAATTGCCATCATCAGGGGGCGCGTTCTGTTCGCGCCGTGAATGAA	1080
1081	ATGTCAGATAACTGGCGACAGGCCTGTTATAAAAATAAACAATACATTATGGGAAAGTAA	1140
1141	TACAGCTGCAGCGTTTCTGAACAGAAAGTCACAGTTTTTATATACAACGGGTAAATAAA	1200
1201	${\tt GAGTTAAGCATGAAGAAGATGTTTATGGCGGTTTTATTTGCATTAGCTTCTGTTAATGCAA}$	1260
1261	ATGGCGGCGGATTGTGCTAAAGGTAAAATTGAGTTTTCCAAGTATAATGAGGATGACACA	1320
1321	TTTACAGTGAAGGTTGACGGGAAAGAATACTGGACCAGTCGCTGGAATCTGCAACCGTTA	1380
1381	CTGCAAAGTGCTCAGTTGACAGGAATGACTGTCACAATCAAATCCAGTACCTGTGAATCA	1440
1441	GGCTCCGGATTTGCTGAAGTGCAGTTTAATAATGACTGA 1479 FIG.	5
	rig.	J

1	gcggatccAA	GGAATTTACC	T'I'AGAC'I'I'C'I'	CGACTGCAAA	GACGTATGTA
51	GATTCGCTGA	ATGTCATTCG	CTCTGCAATA	GGTACTCCAT	TACAGACTAT
101	TTCATCAGGA	GGTACGTCTT	TACTGATGAT	TGATAGTGGC	TCAGGGGATA
151	ATTTGTTTGC	AGTTGATGTC	AGAGGGATAG	ATCCAGAGGA	AGGGCGGTTT
201	AATAATCTAC	GGCTTATTGT	TGAACGAAAT	AATTTATATG	TGACAGGATT
251	TGTTAACAGG	ACAAATAATG	TTTTTTATCG	CTTTGCTGAT	TTTTCACATG
301	TTACCTTTCC	AGGTACAACA	GCGGTTACAT	TGTCTGGTGA	CAGTAGCTAT
351	ACCACGTTAC	AGCGTGTTGC	AGGGATCAGT	CGTACGGGGA	TGCAGATAAA
401	TCGCCATTCG	TTGACTACTT	CTTATCTGGA	TTTCCTGTCG	CATAGTGGAA
451	CCTCACTGAC	GCAGTCTGTG	GCAAGAGCGA	TGTTACGGTT	TGTTACTGTG
501	ACAGCTGAAG	CTTTACGTTT	TCGGCAAATA	CAGAGGGGAT	TTCGTACAAC
551	ACTGGATGAT	CTCAGTGGGC	GTTCTTATGT	AATGACTGCT	GAAGATGTTG
601	ATGTTACATT	GAACTGGGGA	AGGTTGAGTA	GCGTCCTGCC	TGACTATCAT
651	GGACAAGACT	CTGTTCGTGT	AGGAAGAATT	TCTTTTGGAA	GCATTAATGC
701	AATTCTGGGA	AGCGTGGCAT	TAATACTGAA	TTGTCATCAT	CATGCATCGC
751	GAGTTGCCAG	AATGGCATCT	GATGAGTTTC	CTTCTATGTG	TCCGGCAGAT
801	GGAAGAGTCC	GTGGGATTAC	GCACAATAAA	ATATTGTGGG	ATTCATCCAC
851	TCTGGGGGCA	ATTCTGATGC	GCAGAACTAT	ATATTGTGGG	ATTCATCCAC
901	TGAAAAAAAC	ATTATTAATA	GCTGCATCGC	TTTCATTTT	TTCAGCAAGT
951	GCGCTGGCGA	CGCCTGATTG	TGTAACTGGA	AAGGTGGAGT	ATACAAAATA
1001	TAATGATGAC	GATACCTTTA	CAGTTAAAGT	GGGTGATAAA	. GAATTATTTA
1051	CCAACAGATG	GAATCTTCAG	TCTCTTCTTC	TCAGTGCGCA	AATTACGGGG
1101	ATGACTGTAA	CCATTAAAAC	TAATGCCTGT	CATAATGGAG	GGGGATTCAC
1151	CGAAGTTATT	TTTCGTTGAC	TCAGAATAGC	TgCAGTGAAA	AT

FIG. 6A

7/19

1	gcggatccga	tgacgatgac	aaaAAGGAAT	TTACCTTAGA	CTTCTCGACT
51	GCAAAGACGT	ATGTAGATTC	GCTGAATGTC	ATTCGCTCTG	CAATAGGTAG
101	TCCATTACAG	ACTATTTCAT	CAGGAGGTAC	GTCTTTACTG	ATGATTGAT
151	GTGGCTCAGG	GGATAATTTG	TTTGCAGTTG	ATGTCAGAGG	GATAGATCC
201	GAGGAAGGGC	GGTTTAATAA	TCTACGGCTT	ATTGTTGAAC	GAAATAATTI
251	ATATGTGACA	GGATTTGTTA	ACAGGACAAA	TAATGTTTTT	TATCGCTTTC
301	CTGATTTTTC	ACATGTTACC	TTTCCAGGTA	CAACAGCGGT	TACATTGTCT
351	GGTGACAGTA	GCTATACCAC	GTTACAGCGT	GTTGCAGGGA	TCAGTCGTAC
401	GGGGATGCAG	ATAAATCGCC	ATTCGTTGAC	TACTTCTTAT	CTGGATTTAA
451	TGTCGCATAG	TGGAACCTCA	CTGACGCAGT	CTGTGGCAAG	AGCGATGTTA
501	CGGTTTGTTA	CTGTGACAGC	TGAAGCTTTA	CGTTTTCGGC	AAATACAGAC
551	GGGATTTCGT	ACAACACTGG	ATGATCTCAG	TGGGCGTTCT	TATGTAATGA
601	CTGCTGAAGA	TGTTGATCTT	ACATTGAACT	GGGGAAGGTT	GAGTAGCGTC
651	CTGCCTGACT	ATCATGGACA	AGACTCTGTT	CGTGTAGGAA	GAATTTCTTT
701	TGGAAGCATT	AATGCAATTC	TGGGAAGCGT	GGCATTAATA	CTGAATTGTC
751	ATCATCATGC	ATCGCGAGTT	GCCAGAATGG	CATCTGATGA	GTTTCCTTCT
801	ATGTGTCCGG	CAGATGGAAG	AGTCCGTGGG	ATTACGCACA	ATAAAATATI
851	GTGGGATTCA	TCCACTCTGG	GGGCAATTCT	GATGCGCAGA	ACTATTAGCA
901	GTTGAGGGGG	TAAAATGAAA	AAAACATTAT	TAATAGCTGC	ATCGCTTTCA
951	TTTTTTCAG	CAAGTGCGCT	GGCGACGCCT	GATTGTGTAA	CTGGAAAGGT
1001	GGAGTATACA	AAATATAATG	ATGACGATAC	CTTTACAGTT	AAAGTGGGTG
1051	ATAAAGAATT	ATTTACCAAC	AGATGGAATC	TTCAGTCTCT	TCTTCTCAGT
1101	GÇGCAAATTA	CGGGGATGAC	TGTAACCATT	AAAACTAATG	CCTGTCATA
1151	TGGAGGGGGA	TTCAGCGAAG	TTATTTTTCG	TTGACTCAGA	ATAGCTGCAG
1201	TGAAAAT				



1	gcatatgcat	caccatcacc	atcacAAGGA	ATTTACCTTA	GACT TCTCGA
51	CTGCAAAGAC	GTATGTAGAT	TCGCTGAATG	TCATTCGCTC	TGCAATAGGT
101	ACTCCATTAC	AGACTATTTC	ATCAGGAGGT	ACGTCTTTAC	TGATGATTGA
151	TAGTGGCTCA	GGGGATAATT	TGTTTGCAGT	TGATGTCAGA	GGGATAGATC
201	CAGAGGAAGG	GCGGTTTAAT	AATCTACGGC	TTATTGTTGA	ACGAAATAAT
251	TTATATGTGA	CAGGATTTGT	TAACAGGACA	AATAATGTTT	TTTATCGCTT
301	TGCTGATTTT	TCACATGTTA	CCTTTCCAGG	TACAACAGCG	GTTACATTGT
351	CTGGTGACAG	TAGCTATACC	ACGTTACAGC	GTGTTGCAGG	GATCAGTCGT
401	ACGGGGATGC	AGATAAATCG	CCATTCGTTG	ACTACTTCTT	ATCTGGATTT
451	AATGTCGCAT	AGTGGAACCT	CACTGACGCA	GTCTGTGGCA	AGAGCGATGT
501	TACGGTTTGT	TACTGTGACA	GCTGAAGCTT	TACGTTTTCG	GCAAATACAG
551	AGGGGATTTC	GTACAACACT	GGATGATCTC	AGTGGGCGTT	CTTATGTAAT
601	GACTGCTGAA	GATGTTGATC	TTACATTGAA	CTGGGGAAGG	TTGAGTAGCG
651	TCCTGCCTGA	CTATCATGGA	CAAGACTCTG	TTCGTGTAGG	AAGAATTTCT
701	TTTGGAAGCA	TTAATGCAAT	TCTGGGAAGC	GTGGCATTAA	TACTGAATTG
751	TCATCATCAT	GCATCGCGAG	TTGCCAGAAT	GGCATCTGAT	GAGTTTCCTT
801	CTATGTGTCC	GGCAGATGGA	AGAGTCCGTG	GGATTACGCA	СААТААААТА
851	TTGTGGGATT	CATCCACTCT	GGGGGCAATT	CTGATGCGCA	GAACTATTAG
901	CAGTTGAGGG	GGTAAAATGA	AAAAAACATT	ATTAATAGCT	GCATCGCTTT
951	CATTTTTTC	AGCAAGTGCG	CTGGCGACGC	CTGATTGTGT	AACTGGAAAG
1001	GTGGAGTATA	СААААТАТАА	AGATGACGAT	ACCTTTACAG	TTAAAGTGGG
1051	TGATAAAGAA	ТТАТТТАССА	ACAGATGGAA	TCTTCAGTCT	CTTCTTCTCA
1101	GTGCGCAAAT	TACGGGGATG	ACTGTAACCA	TTAAAACTAA	TGCCTGTCAT
1151	AATGGAGGGG	GATTCAGCGA	AGTTATTTT	CGTTGACTCA	GAATAGCTC
1201	GTGAAAATAG	CAGGCGGAGA	TaTCgATAAA	TGTTA	

FIG. 6C SUBSTITUTE SHEET (RULE 26)

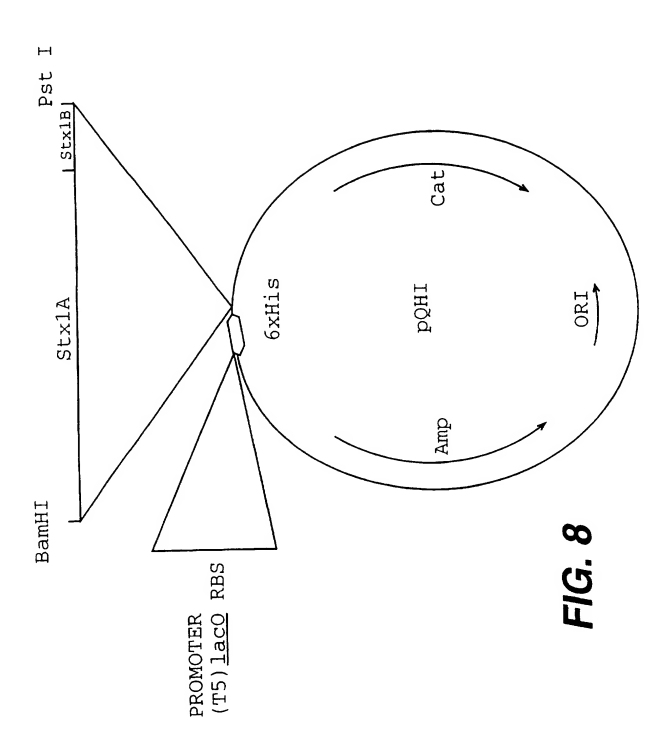
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51	CTTCGTTAAA	TAGTATACGG	ACAGAGATAT	CGACCCCTCT	TGAACATATA
101	TCTCAGGGGA	CCACATCGGT	GTCTGTTATT	AACCACACCC	CACCGGGCAG
151	TTATTTTGCT	GTGGATATAC	GAGGGCTTGA	TGTCTATCAG	GCGCGTTTTG
201	ACCATCTTCG	TCTGATTATT	GAGCAAAATA	ATTTATATGT	GGCCGGGTTC
251	GTTAATACGG	CAACAAATAC	TTTCTACCGT	TTTTCAGATT	TTACACATAT
301	ATCAGTGCCC	GGTGTGACAA	CGGTTTCCAT	GACAACGGAC	AGCAGTTATA
351	CCACTCTGCA	ACGTGTCGCA	GCGCTGGAAC	GTTCCGGAAT	GCAAATCAGT
401	CGTCACTCAC	TGGTTTCATC	ATATCTGGCG	TTAATGGAGT	TCAGTGGTAA
451	TACAATGACC	AGAGATGCAT	CCAGAGCAGT	TCTGCGTTTT	GTCACTGTCA
501	CAGCAGAAGC	CTTACGCTTC	AGGCAGATAC	AGAGAGAATT	TCGTCAGGCA
551	CTGTCTGAAA	CTGCTCCTGT	GTATACGATG	ACGCCGGGAG	ACGTGGACCT
601	CACTCTGAAC	TGGGGGCGAA	TCAGCAATGT	GCTTCCGGAG	TATCGGGGAG
651	AGGATGGTGT	CAGAGTGGGG	AGAATATCCT	ТТААТААТАТ	ATCAGCGATA
701	CTGGGGACTG	TGGCCGTTAT	ACTGAATTGC	CATCATCAGG	GGGCGCGTTC
751	TGTTCGCGCC	GTGAATGAAG	AGAGTCAACC	AGAATGTCAG	ATAACTGGCG
801	ACAGGCCTGT	ТАТАААААТА	AACAATACAT	TATGGGAAAG	TAATACAGCT
851	GCAGCGTTTC	TGAACAGAAA	GTCACAGTTT	ТТАТАТАСАА	CGGGTAAATA
901	AAGGAGTTAA	GCATGAAGAA	GATGTTTATG	GCGGTTTTAT	TTGCATTAGC
951	TTCTGTTAAT	GCAATGGCGG	CGGATTGTGC	TAAAGGTAAA	ATTGAGTTTT
.001	CCAAGTATAA	TGAGGATGAC	ACATTTACAG	TGAAGGTTGA	CGGGAAAGAA
.051	TACTGGACCA	GTCGCTGGAA	TCTGCAACCG	TTACTGCAAA	GTGCTCAGTT
101	GACAGGAATG	ACTGTCACAA	TCAAATCCAG	TACCTGTGAA	TCAGGCTCCG
151	GATTTGCTGA	AGTGCAGTTT	AATAATGACT	GAGGCATAAg	CTTATTCGTG
1201	G				

FIG. 7A SUBSTITUTE SHEET (RULE 26)

1	gcggatccga	tgacgatgac	aaaCGGGAGT	TTACGATAGA	CTTTTCGACC
51	CAACAAAGTT	ATGTCTCTTC	GTTAAATAGT	ATACGGACAG	AGATATCGAC
101	CCCTCTTGAA	CATATATCTC	AGGGGACCAC	ATCGGTGTCT	GTTATTAACC
151	ACACCCCACC	GGGCAGTTAT	TTTGCTGTGG	ATATACGAGG	GCTTGATGTC
201	TATCAGGCGC	GTTTTGACCA	TCTTCGTCTG	ATTATTGAGC	AAAATAATTT
251	ATATGTGGCC	GGGTTCGTTA	ATACGGCAAC	AAATACTTTC	TACCGTTTTT
301	CAGATTTTAC	ACATATATCA	GTGCCCGGTG	TGACAACGGT	TTCCATGACA
351	ACGGACAGCA	GTTATACCAC	TCTGCAACGT	GTCGCAGCGC	TGGAACGTTC
401	CGGAATGCAA	ATCAGTCGTC	ACTCACTGGT	TTCATCATAT	CTGGCGTTAA
451	TGGAGTTCAG	TGGTAATACA	ATGACCAGAG	ATGCATCCAG	AGCAGTTCTG
501	CGTTTTGTCA	CTGTCACAGC	AGAAGCCTTA	CGCTTCAGGC	AGATACAGAG
551	AGAATTTCGT	CAGGCACTGT	CTGAAACTGC	TCCTGTGTAT	ACGATGACGC
601	CGGGAGACGT	GGACCTCACT	CTGAACTGGG	GGCGAATCAG	CAATGTGCTT
651	CCGGAGTATC	GGGGAGAGGA	TGGTGTCAGA	GTGGGGAGAA	TATCCTTTAA
701	TAATATATCA	GCGATACTGG	GGACTGTGGC	CGTTATACTG	AATTGCCATC
751	ATCAGGGGGC	GCGTTCTGTT	CGCGCCGTGA	ATGAAGAGAG	TCAACCAGAA
801	TGTCAGATAA	CTGGCGACAG	GCCTGTTATA	AAAATAAACA	ATACATTATG
851	GGAAAGTAAT	ACAGCTGCAG	CGTTTCTGAA	CAGAAAGTCA	CAGTTTTTAT
901	ATACAACGGG	TAAATAAAGG	AGTTAAGCAT	GAAGAAGATG	TTTATGGCGG
951	TTTTATTTGC	ATTAGCTTCT	GTTAATGCAA	TGGCGGCGGA	TTGTGCTAAA
1001	GGTAAAATTG	AGTTTTCCAA	GTATAATGAG	GATGACACAT	TTACAGTGAA
1051	GGTTGACGGG	AAAGAATACT	GGACCAGTCG	CTGGAATCTG	CAACCGTTAC
1101	TGCAAAGTGC	TCAGTTGACA	GGAATGACTG	TCACAATCAA	ATCCAGTACC
1151	TGTGAATCAG	GCTCCGGATT	TGCTGAAGTG	CAGTTTAATA	ATGACTGAGG
1201	CATAAgCTtA	TTCGTGG			

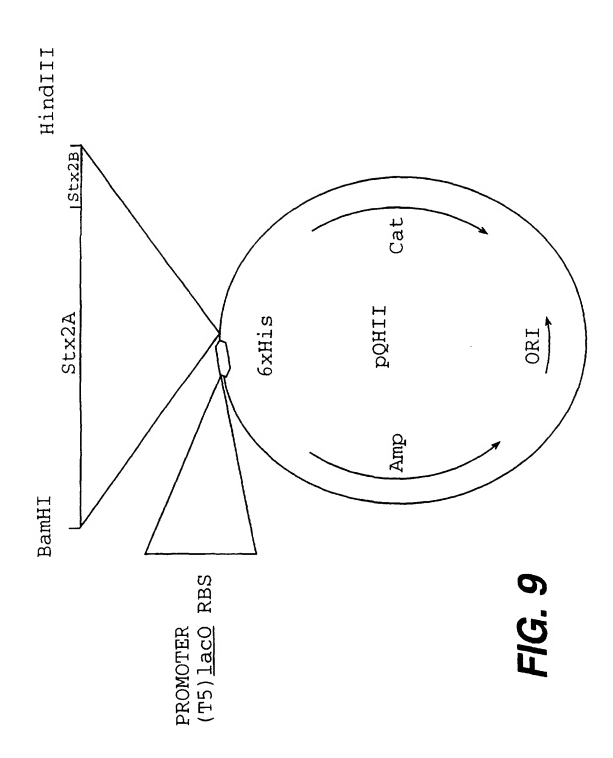


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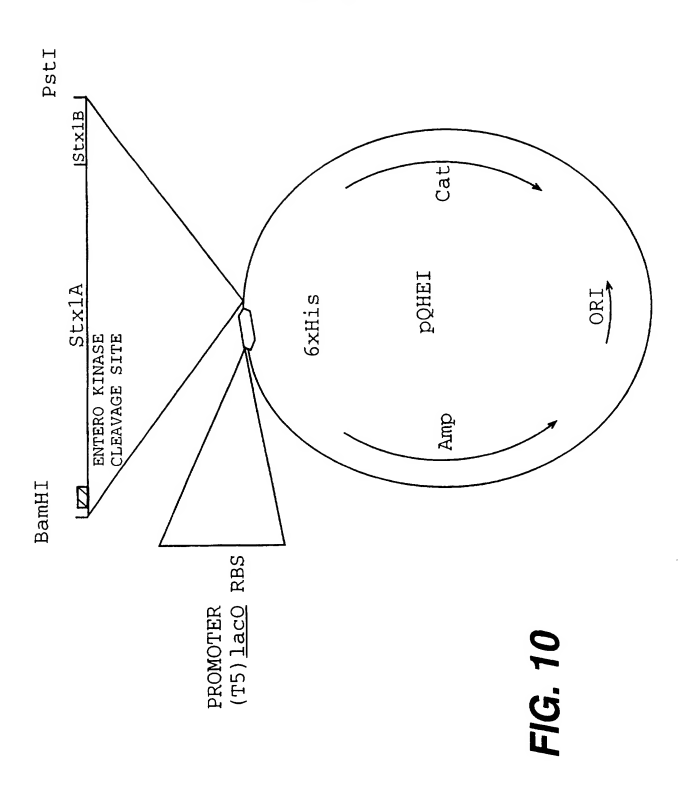
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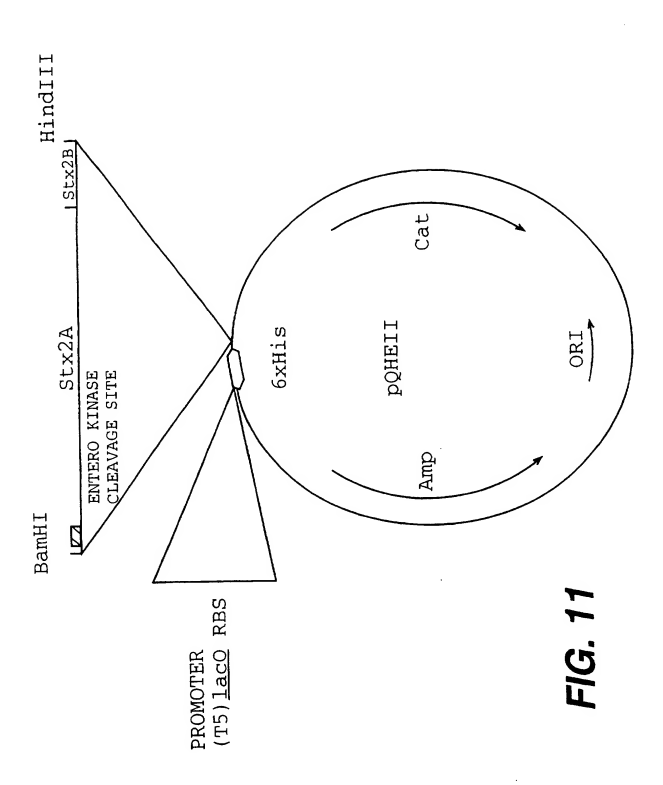


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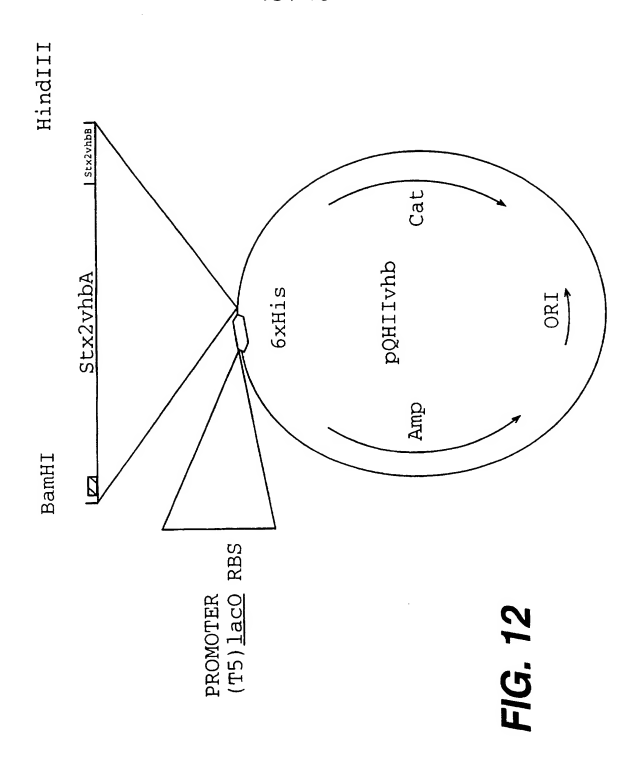


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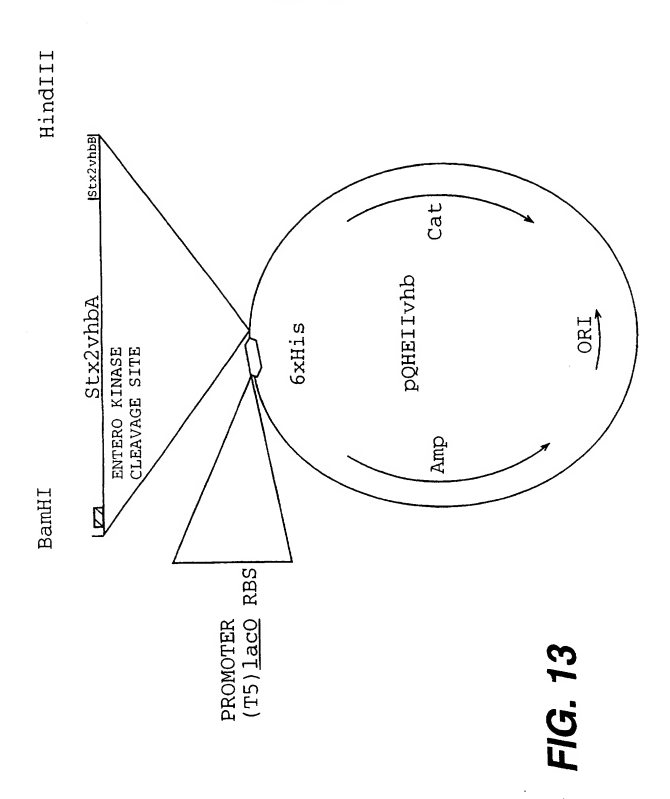


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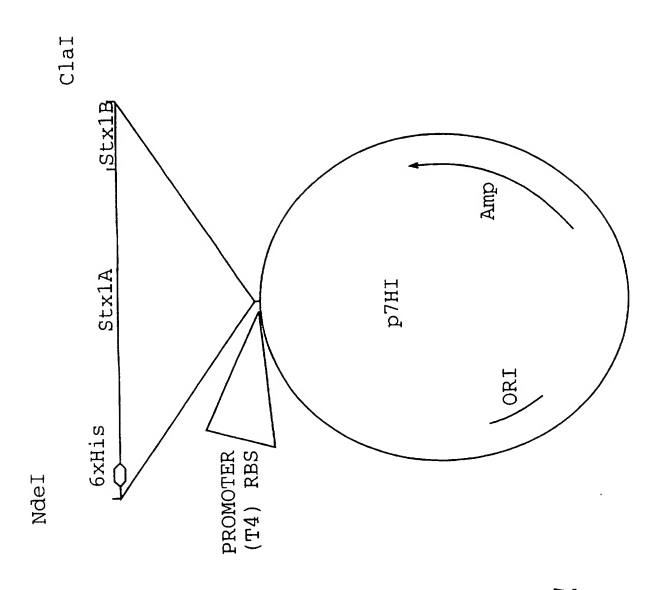
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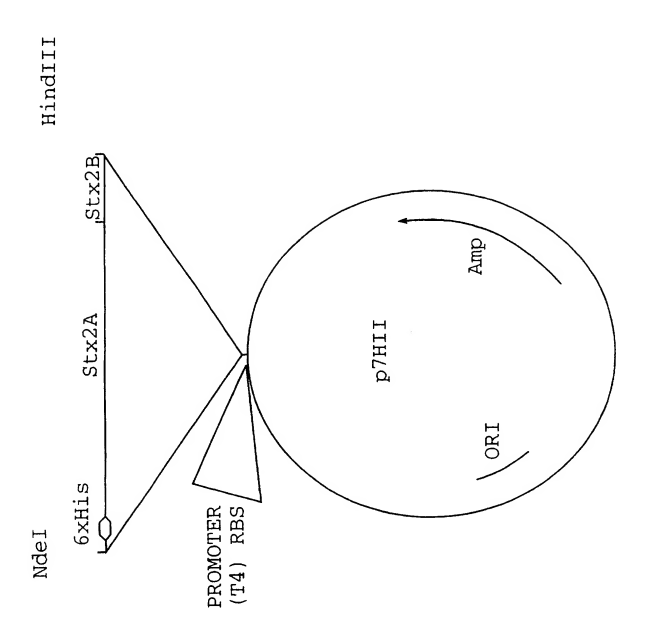
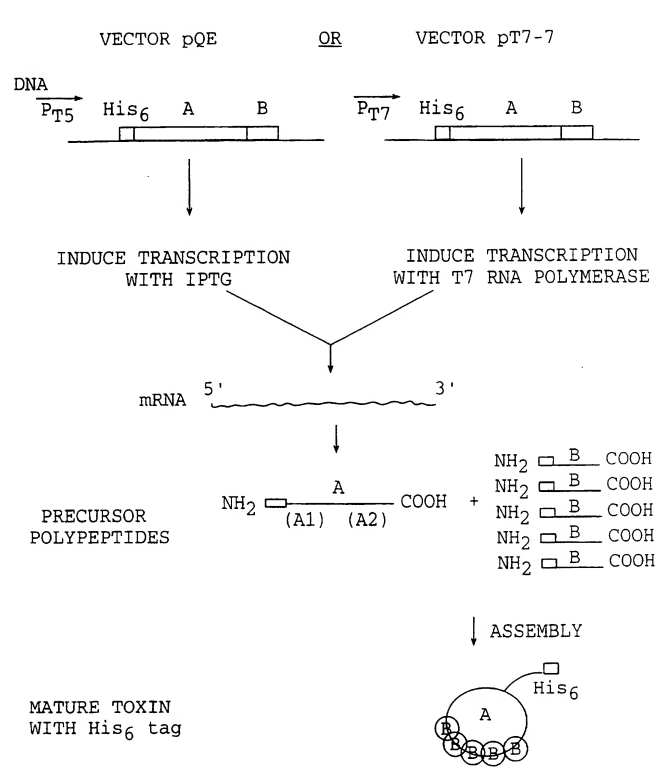


FIG. 15

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P, PROMOTER. . 6 HISTIDINE RESIDUES.

FIG. 16
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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:		(11) International Publication Number: WO 98/11229
C12N 15/31, 15/62, C07K 14/245, 14/25, 16/12, A61K 39/40, 48/00	A3	(43) International Publication Date: 19 March 1998 (19.03.98)
(21) International Application Number: PCT/US9 (22) International Filing Date: 9 September 1997 (0)		CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
 (30) Priority Data: 60/025,637 10 September 1996 (10.09.9) (71) Applicant: HENRY M. JACKSON FOUNDATION F ADVANCEMENT OF MILITARY MEDICINE (Suite 600, 1401 Rockville Pike, Rockville, MD 208 (72) Inventors: O'BRIEN, Alison, D.; 5514 Charlcon Bethesda, MD 20817 (US). SCHMITT, Clare, I Glasgow Drive, Rockville, MD 20853 (US). (74) Agents: GARRETT, Arthur, S. et al.; Finnegan, He Farabow, Garrett & Dunner, L.L.P., 1300 I Street Washington, DC 20005-3315 (US). 	FOR THE (US/US) 352 (US) te Roak K.; 46	Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments. [88] Date of publication of the international search report: 16 April 1998 (16.04.98)

(54) Title: HISTIDINE-TAGGED SHIGA TOXINS, TOXOIDS, AND PROTEIN FUSIONS WITH SUCH TOXINS AND TOXOIDS, METHODS FOR THE PURIFICATION AND PREPARATION THEREOF

(57) Abstract

The present invention describes the isolation and purification of biologically and immunologically active histidine-tagged Shiga toxins (His-tagged), a toxin associated with HC and the potentially life-threatening sequela HUS transmitted by strains of pathogenic bacteria. The present invention describes how his-tagging greatly simplifies and expedites purifying Shiga toxins, and describes an improved method for such purification. One aspect of the invention is obtaining and using Shiga toxids that are immunoreactive but not toxic. Another aspect of the invention is obtaining and using fusion proteins of His-tagged Shiga toxins or toxoids. Yet another aspect of the invention is obtaining and using antibodies to His-gagged Shiga toxins, toxoids, or Shiga toxin/toxoid fusion proteins.

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According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) IPC 6 C12N C07K A61K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

C. DOCUM	ENTS CONSIDERED TO BE RELEVANT	
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	WO 96 12802 A (OPHIDIAN PHARM INC; WILLIAMS JAMES A (US); PADHYE NISHA V (US); KI) 2 May 1996 see page 24, line 12 - line 17 see page 25, line 20 - line 30 see page 26, line 1 - line 16 see page 28, line 9 - line 17 see page 30, line 20 - page 31, line 5	1-7
Υ .	GORDON V.M. ET AL.: "An enzymatic mutant of Shiga-like toxin II variant is a vaccine candidate for Edema disease of swine." INFECTION AND IMMUNITY, vol. 60, no. 2, 1992, pages 485-490, XP002054488 cited in the application see the whole document	1-7
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Date of the actual completion of the international search 4 February 1998	Date of mailing of the international search report 0 5. 03. 98
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, Fax: (+31-70) 340-3016	Authorized officer Mandl, B

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Further documents are listed in the continuation of box C.

Patent family members are listed in annex.

1

		PC1/03 97/13830
	lation) DOCUMENTS CONSIDERED TO BE RELEVANT	
ategory °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	HOCHULI E. ET AL.: "Genetic approach to facilitate purification of recombinant proteins with a novel metal chelate adsorbent." BIOTECHNOLOGY, vol. 6, no. 11, 1988, pages 1321-1325, XP002054489 see the whole document	1-4
Ą	LINDGREN S. W. ET AL.: "The specific activities of Shiga-like toxin type II (SLT-II) and SLT-II-related toxin of Enterohemorrhagic Escherichia coli differ when measured by Vero cell cytotoxicity but not by mouse lethality." INFECTION AND IMMUNITY, vol. 62, no. 2, 1994, pages 623-631, XP002054490 cited in the application see the whole document, especially Table 1	1-8
A	LI J.: "Bacterial toxins." CURRENT OPINION IN STRUCTURAL BIOLOGY, vol. 2, no. 4, 1992, pages 545-556, XP000335477 see page 550, left-hand column, paragraph 4 - right-hand column, paragraph 1	1-8
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Form PCT/ISA/210 (continuation of second sheet) (July 1992)

INTERNATIONAL SEARCH REPORT

Int....ational application No. PCT/US 97/15836

Box	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This Inte	ernational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. X	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely: see FURTHER INFORMATION sheet PCT/ISA/210
2.	Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
3.	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rute 6.4(a).
BxII	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This into	emational Searching Authority found multiple inventions in this international application, as follows:
1.	As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2.	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3.	As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4.	No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remari	The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

Form PCT/ISA/210 (continuation of first sheet (1)) (July 1992)

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210
Remark: Although claims 5 and 6 are directed to a method of treatment of the human/animal body, the search has been carried out and based on the alleged effects of the compound/composition.

information on patent family members

PCT/US 97/15836

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